<b>Patient Identification</b>	(record all	l dates as m	nm/dd/yyyy	<b>(</b> )								
*First Name		*Middle Name				*Last Name			Last Name Soundex			
Alternate Name Type (ex: Alias, Married)		*First Name			*Middle Name		,	*Last Name				
	e 🗆 Homele	ss   Military	•	*Currer	t Addres	ss, Street				Address Date		
	Shelter   Temporary  City		County		State/Country			*ZIP Code				
*Medical Record Number			*Other ID Type			*Number						
U.S. Department of Health and Human Services	(Pat					ase Repor			С	Centers for Disease Control and Prevention (CDC)		
Health Department U		ecord all da	ites as mr	n/dd/yyyy)	)	Form	n approv	ed OMB	no. 0920	)-0573 Exp. 11/30/2022		
Date Received at Health Department			eHARS D	eHARS Document UID			State Number					
Reporting Health Dept—Cit	y/County		City/County Number									
Document Source				Surveillance Method  Active Passive Follow up Reabstraction Unknown								
Did this report initiate a new case investigation?  ☐ Yes ☐ No ☐ Unknown				Report Medium  ☐ 1-Field visit ☐ 2-Mailed ☐ 3-Faxed ☐ 4-Phone ☐ 5-					-Electronic transfer    6-CD/disk			
Facility Providing Info	rmation (	record all d	ates as m	m/dd/yyy	<b>/</b> )							
Facility Name								*Phone				
*Street Address												
City	Coun	ty			State/0	Country		*ZIP Cod	de			
Facility     Inpatient:     Outpatient:     □ P       Type     □ Hospital     □ Adult HIV clini       □ Other, specify     □ Other, specify			ic CTS			□ STD clinic □			□ Labora	her Facility: ☐ Emergency room  Laboratory ☐ Corrections ☐ Unknown  Other, specify		
			Person Completing Form				*Phone					
Patient Demographic	s (record a	ıll dates as	mm/dd/yy	уу)								
Sex Assigned at Birth  ☐ Male ☐ Female ☐ Unk	nown			ntry of Birth		idency (please s	necify)					
Date of Birth /	1					Date of Birth	/	/				
Vital Status □ 1-Alive □ 2	 2-Dead	С	ate of Deat	h /	/		State o	of Death				
			-	ale-to-fema	le (MTF)	□ Transgender	r female-	to-male (	FTM) 🗆	Unknown		
□ Additional gender identity (specify)  Ethnicity □ Hispanic/Latino □ Not Hispanic/Latino □ Unknown												
Race □ American Indian/Alaska Native □ Asian □ Black/African American (check all that apply) □ Native Hawaiian/Other Pacific Islander □ White □ Unknown												
Residence at Diagnos	is (add ad	lditional add	dresses in	Commen	ts) (rec	ord all dates	as mm/	dd/yyyy	·)			
Address Event Type (check all that apply to address	ss below) $\Box$	Residence at	HIV diagnos	sis □ Resi	dence at	stage 3 (AIDS) dia	agnosis	□ Checl	k if <u>SAM</u>	as current address		
Address Type   Residentia												
*Street Address												
City	Cour	nty		(	State/Cou	ate/Country *ZIP Code				*ZIP Code		

Public reporting burden of this collection of information is estimated to average 20 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to CDC, Project Clearance Officer, 1600 Clifton Road, MS D-74, Atlanta, GA 30333, ATTN: PRA (0920-0573). **Do not send the completed form to this address.** 

STATE/LOCAL USE	ONLY											
*Provider Name (Last, F	_						*Phone	e (	)			
Hospital/Facility	. ,							`	,			
Facility of Diagnosi				(1150)	01 1 1 0 0 0 0 0							
Diagnosis Type (check a	Il that apply to	tacility belov	w) □ HIV □ Stage 3	(AIDS)	□ Check if <u>SAME</u>				ion			
Facility Name						*Pho	one (	)				
*Street Address												
City		•					*ZIP C					
Facility Type <u>Inpatient:</u> □ Other, s	☐ Hospital specify	☐ Adult HIV		<i>Screenin</i> □ CTS	Other Facility: ☐ Emergency room ☐ Laboratory ☐ Corrections ☐ Unknown							
		☐ Other, spe	ecify	□ Other,	specify		□ Other	, speci	fy			
*Provider Name			*Provider Phone ( )	)		Spe	cialty					
Dationt History (ros	nand to all	auestions	) (record all dates as ।	mm/dd/v	<b>2004)</b>	- Dodiatri	c Diek	(plac	aco on	tor i	n Comment	
			sis of HIV infection, this p			rediatii	CINISK	(pie	ase em	er ii	Comment	
Sex with male			р					□ Yes	. □ No		Unknown	
Sex with female								□ Yes			Unknown	
	rugo.							□ Yes			Unknown	
Injected nonprescription dr Received clotting factor for		occulation d	ligardar					□ Yes			Unknown	
Specify clotting factor:	петторина/с	oagulalion u	isorder	Date	received /	/		⊔ res	i □ No	, ⊔	Olikilowii	
HETEROSEXUAL relation	ns with any o	of the follow	ring:									
HETEROSEXUAL contact	with intraven	ous/injection	drug user					□ Yes	. □ No		Unknown	
HETEROSEXUAL contact with hisexual male								□ Yes	□ No		Unknown	
HETEROSEXUAL contact with person with hemophilia/coagulation disorder with documented HIV infection								□ Yes	□ No		Unknown	
								Unknown				
	· · · · · · · · · · · · · · · · · · ·									Unknown		
								Unknown				
									Unknown			
			t date received /			<b>5</b> )		00		, _	O I II I I I I I I I I I I I I I I I I	
Received transplant of tiss					<u> </u>			□ Yes	. □ No	, ,	Unknown	
Worked in a healthcare or			- Initiation					□ Yes			Unknown	
			sidered					□ 103		, ⊔	Offictiowit	
If occupational exposure is being investigated or considered as primary mode of exposure, specify occupation and setting:												
Other documented risk (please include detail in Comments)								□ Yes	□ No	) 🗆	Unknown	
Clinical, Acuta HIV	Infoction	and Onn	ortunistic Illnesses	(recerd	all dates as my							
			o items below; enter document				ection an	d	⊓ Vac	□ Nc	o 🗆 Unknowi	
enter patient or provider report	of previous ne	gative HIV test	t in HIV Testing History section	1.								
Clinical signs/symptoms consistent with acute retroviral syndrome (e.g., fever, malaise/fatigue, myalgia, pharyngitis, rash, lymphadenopathy)? Date of sign/symptom onset//							n,		□ Yes	□ No	o 🗆 Unknowi	
Other evidence suggestive Date of evidence/_	e of acute HIV	/ infection?	If YES, please describe:						□ Yes	□ No	o □ Unknowi	
Opportunistic Illnesses		_										
Diagnosis	Dx Da	ate	Diagnosis		Dx Date	Diagnosis		-		Dx D	)ate	
Candidiasis, bronchi, trachea, or	lungs		Herpes simplex: chronic ulcers duration), bronchitis, pneumon esophagitis			M. tuberculos	is, pulmona	ary'				
Candidiasis, esophageal			Histoplasmosis, disseminated	or		M. tuberculosi		ated or				
Carcinoma, invasive cervical			extrapulmonary extrapulmonary extrapulmonary lososporiasis, chronic intestinal (>1 mo. Mycobacterium, of oth					er/unidentified				
Coccidioidomycosis, disseminated	d or		duration)  Kaposi's sarcoma			species, disse			ulmonary	$\vdash$		
extrapulmonary	u Ui		naposis salloulla			neumocystis	, priedilloll	ia		L		
Cryptococcosis, extrapulmonary	nol (s.1		Lymphoma, Burkitt's (or equiva			Progressive p		12 mo	. period	<u> </u>		
Cryptosporidiosis, chronic intestinal (>1 Lymphoma, immunoblastic (or equivalent) Progressive multifocal leukoencephalopathy						'						
Cytomegalovirus disease (other tl iver, spleen, or nodes)	han in		Lymphoma, primary in brain			Salmonella se	epticemia, i	ecurrer	nt			
Cytomegalovirus retinitis (with los	ss of		Mycobacterium avium complex			Toxoplasmosi	s of brain, o	onset at	>1 mo.			
vision) HIV encephalopathy			kansasii, disseminated or extra	pulmonary		of age Wasting syndr	rome due to	o HIV				
116 - 41	oithar tubareule	oio dioanos!!	novo provido DVCT Coos Numbe			J = J - L						

Laboratory Data (record additional tests and tests not specified	l below in Comments) (record all dates as mm/dd/yyyy)						
HIV Immunoassays (Nondifferentiating)							
TEST 1   HIV-1 IA   HIV-1/2 IA   HIV-1/2 Ag/Ab   HIV-1 WB   HIV-1 II							
Test brand name/Manufacturer							
Facility name							
Result □ Positive □ Negative □ Indeterminate	Collection Date/ Point-of-care rapid test						
TEST 2   HIV-1 IA   HIV-1/2 IA   HIV-1/2 Ag/Ab   HIV-1 WB   HIV-1 II							
Test brand name/Manufacturer							
	Provider name						
Result □ Positive □ Negative □ Indeterminate	Collection Date/ Point-of-care rapid test						
HIV Immunoassays (Differentiating)							
☐ HIV-1/2 type-differentiating immunoassay (differentiates between HIV-1 Ab and HIV-2 Ab)	Role of test in diagnostic algorithm  ☐ Screening/initial test ☐ Confirmatory/supplemental test						
Test brand name/Manufacturer							
Facility name							
HIV-1 indeterminate ☐ HIV-2 indeterminate							
	Collection Date// Doint-of-care rapid test						
	<sup>1</sup> Always complete the overall interpretation. Complete the analyte results when available.						
☐ HIV-1/2 Ag/Ab differentiating immunoassay (differentiates between HIV Ag							
Test brand name/Manufacturer	· · · · · · · · · · · · · · · · · · ·						
Facility name	Provider name						
Result □ Ag positive □ Ab positive □ Both (Ag and Ab positive) □ Negative							
Collection Date// Point-of-care rapid test	e invalid						
□ HIV-1/2 Ag/Ab and type-differentiating immunoassay (differentiates amon	α HIV-1 Aα HIV-1 Ab and HIV-2 Ab)						
Test brand name/Manufacturer							
Facility name	Provider name						
Result <sup>2</sup> Overall interpretation: □ Reactive □ Nonreactive Index value							
Analyte results: HIV-1 Ag: □ Reactive □ Nonreactive □ Not report							
HIV-1 Ab:  Reactive  Nonreactive  Reactive  Reactive							
HIV-2 Ab: □ Reactive □ Nonreactive □ Reactive □							
Collection Date// Deactive Date/20							
HIV Detection Tests (Qualitative)	Complete the overall interpretation and the analyte results.						
TEST □ HIV-1 RNA/DNA NAAT (Qualitative) □ HIV-1 culture □ HIV-2 RNA/I	DNA NAAT (Qualitative)   HIV-2 culture						
Test brand name/Manufacturer							
Facility name							
Result □ Positive □ Negative □ Indeterminate	Collection Date//						
HIV Detection Tests (Quantitative viral load) Note: Include earliest test a							
TEST 1   HIV-1 RNA/DNA NAAT (Quantitative viral load)   HIV-2 RNA/DNA							
Test brand name/Manufacturer							
Facility name							
Result □ Detectable □ Undetectable Copies/mL	Log Collection Date / /						
TEST 2   HIV-1 RNA/DNA NAAT (Quantitative viral load)   HIV-2 RNA/DNA							
Test brand name/Manufacturer							
Facility name	Provider name						
Result   Detectable Undetectable Copies/mL	LogCollection Date / / /						
Drug Resistance Tests (Genotypic)	Conceilon Date						
TEST □ HIV-1 Genotype (Unspecified)	Test brand name/Manufacturer						
Lab name	Facility name						
Provider name	Collection Date / /						
Immunologic Tests (CD4 count and percentage)	Concolion Bate						
	CD4 percentage % Collection Date / /						
Test brand name/Manufacturer							
Facility name							
	Provider name % Collection Date / /						
Test brand name/Manufacturer	Provider name						
	CD4 percentage% Collection Date//						
Test brand name/Manufacturer							
Facility name	Provider name						
Documentation of Tests	add a control of the state of t						
Did documented laboratory test results meet approved HIV diagnostic algority YES, provide specimen collection date of earliest positive test for this algorithm to the above and it is a few and the f	gorithm / /						
Complete the above only if none of the following were positive for <b>HIV-1</b> : Western differentiating immunoassay (supplemental test), stand-alone p24 antigen, or number of the contraction of the contracti	m biot, ir A, culture, viral ibau, qualitative INAA i (HINA of DINA), HIV-1/2 type- icleotide seguence						
If HIV laboratory tests were not documented, is HIV diagnosis documented							
and the state of t	If YES, provide date of diagnosis///						
Date of last documented negative HIV test (before HIV diagnosis date)							

Treatment/Services Referrals (record all dates as mm/dd/yyyy)									
Has this patient been informed of his/her HIV infection?  ☐ Yes ☐ No ☐ Unknown				heir HIV exposure and counseled by B-Patient □ 9-Unknown					
Evidence of receipt of HIV medical care other than laborate				· · · · · · · · · · · · · · · · · · ·					
	e of medical visi	t or prescription	on//_						
For Female Patient	nical au la Maia	madiant arms		Headhia national delivered live how infented					
This patient is receiving or has been referred for gynecologobstetrical services ☐ Yes ☐ No ☐ Unknown	-	-	Unknown	Has this patient delivered live-born infants? □ Yes □ No □ Unknown					
For Children of Patient (record most recent birth in these b	ooxes; record ad	ditional or mu	Iltiple births in Comn	nents)					
*Child's Name				Child's Date of Birth					
Child's Last Name Soundex	Child's	s State Numb	ber						
Facility Name of Birth				*Phone					
(if child was born at home, enter "home birth")				( )					
Facility Type Inpatient: Outpatient: Other Facility: □ Emergency room									
☐ Hospital ☐ Other, specify ☐ Corrections ☐ Unknown ☐ Other specify									
□ Other, specify □ Other, specify □ Other, specify □ *ZIP Code									
City County State/Country									
				,					
Antiretroviral Use History (record all dates as m									
Main source of antiretroviral (ARV) use information (select or □ Patient interview □ Medical record review □ Prov	*	□ NHM&E	□ Other	Date patient reported information					
Ever taken any ARVs?  Yes  No Unknown	nder report	- IVI IIVIQL	□ Other						
If yes, reason for ARV use (select all that apply)									
□ HIV Tx ARV medications	Da	te began		Date of last use / / /					
□ PrEP ARV medications		_							
□ PEP ARV medications									
□ PMTCT ARV medications		=	_// _//						
□ HBV Tx ARV medications			/						
		te began	_''						
□ Other (specify reason)									
ARV medications	Da	te began		Date of last use / /					
HIV Testing History (record all dates as mm/dd/y	ууу)								
Main source of testing history information (select one)				Date patient reported information					
□ Patient interview □ Medical record review □ Provider re	•								
Ever had previous positive HIV test?   Yes   No   Un	Known		of first positive HIV						
Ever had a negative HIV test? □ Yes □ No □ Unknown			negative HIV test (	(if date is from Lab Data section) / /					
Number of negative HIV tests within the 24 months before	the first positiv			Lab Bata scottoriy					
Number of negative fire tests within the 24 months before	the mat positiv	e test	□ OTINTOWIT						
Comments									
*Local/Optional Fields									

This report to CDC is authorized by law (Sections 304 and 306 of the Public Health Service Act, 42 USC 242b and 242k). Response in this case is voluntary for federal government purposes, but may be mandatory under state and local statutes. Your cooperation is necessary for the understanding and control of HIV. Information in CDC's National HIV Surveillance System that would permit identification of any individual on whom a record is maintained is collected with a guarantee that it will be held in confidence, will be used only for the purposes stated in the assurance on file at the local health department, and will not otherwise be disclosed or released without the consent of the individual in accordance with Section 308(d) of the Public Health Service Act (42 USC 242m).